Search Notes

Application/Control No.	Applicant(s)/Patent under Reexamination		
10/756,849	GABBAY, JEFFREY		
Examiner	Art Unit		
Michael G. Bogart	3761		

SEARCHED					
Class	Subclass	Date	Examiner		
604	359 360 367 375 904	3/13/2006	MGB		
424	402 404	3/13/2006	MGB		
442	123	3/13/2006	MGB		

INT	INTERFERENCE SEARCHED					
Class	Subclass	Date	Examiner			
,						
	<u> </u>					
	1		:			

SEARCH NOTES (INCLUDING SEARCH STRATEGY)				
DATE	EXMR			
3/13/2006	MGB			
1 0				
	DATE			

Therefore, it is clear from the specification what the applicant intended as the scope of the invention as stated in claims 10-13 by including the terms "pharmacological" and "subpharmacological". In view of the above, the applicant respectfully traverses this objection.

Reconsideration and withdrawal of this objection is requested.

Response to Prior Art-Based Rejection

The balance of the Official Action (items 8-11) relates to a single prior art-based rejection of alleged obviousness over one of the Giannessi references in combination with one or the other of the Giannessi references in combination with two secondary references. Specifically, the examiner argues that it is obvious to combine the teachings of WO9959957, which discloses a general formula also including ST1326 among many compounds and its hypothetical combination with other compounds, such as biguanides, together with Dagogo-Jack which discloses that metformin and many other biguanides can be used for the treatment of diabetes, to obtain the solution proposed in the present invention.

WO9959957 gives the following pharmacological data for ST1326:

- IC_{50} of inhibition CPT-1 curve in rat liver mitochondria (Table 1 value 0.75 μ M/I);
- 13-hydroxybutyrate and glucose serum concentration in 24 hours-starved rats, after one hour from intraperitoneal treatment, (ST1326 doses 14.5 mg/2 ml/kg)

and does not indicate ST1326 as a most preferred embodiment.

The person skilled in the art would not find any suggestion in WO9959957 to choose ST1326 from the large number of compounds of formula (I) and to specifically combine it with metformin, one of the many possible biguanides.

The data reported in the present application, showing <u>comparative tests</u> between the combination and the two components alone, demonstrate the synergic effect of the combination and thus establishes non-obviousness.

The synergic effect is demonstrated by the results for the combination ST1326 30 mg/kg - metformin 200 mg/kg shown in table 3, table 4, table 5 and table 6 wherein the reduction of glucose is greater than the single components alone.

In particular, table 3 shows that the combination lowers levels of blood glucose in mice in feeding conditions and 15 hours after the last treatment to 287.3 mg/dl whereas metformin alone

does not lower the blood glucose in respect to the control and metformin lowers it to 376.3 mg/dl.

Table 4 shows that the combination lowers the glucose blood levels in mice in post-absorption conditions and 8 hours after the last treatment to 362.7 mg/dl (the control is 517.7 mg/dl); on the contrary, ST1326 does not influence the levels and metformin alone lowers the blood glucose levels to 433.7

Table 5 shows the results for mice in post-absorption conditions and 6 hours after the last treatment: the control is 360.2 mg/dl, the treatment with the claimed combination lowers the blood glucose levels to 269.9 mg/dl whereas ST1326 is inactive and metformin lowers the levels to the value of 337.8.

This data are confirmed by the results shown in table 6.

In view of the above the skilled person would not be encouraged to combine ST1326 with metformin at concentrations lower than those usually employed in pharmacological treatment.

The skilled person would not be suggested to use concentrations of either drug lower than their well known published pharmaceutical doses (ST1326 = 100 mg/kg/day, metformin = 900 mg/kg/day) and to combine the two active agents since the prior art does not suggest that they could have a synergic effect when used in combination.

It is again emphasized that the skilled person to possible arrive at the claimed combination would be forced to combine and experiment with every single compound disclosed in WO9959957 with each of the biguanides known in the art, without any indication of the possible result achieved by the present invention.

Also the combination of WO9959957 and Giannessi 2003 and/or Dagogo-Jack cannot lead the skilled person to the present invention without an undue burden of experimentation. It is strongly underlined that while Dagogo-Jack gives a list of possible combinations with practical guidelines on their implementation, there no suggestion on the combination of a biguanide with a CPT-I inhibitor.

Therefore, among the many possibilities provided by the prior art, there is no teaching or suggestion of the specific combination of a single, specific biguanides, metformin, with a single specific CPT-I.

The evidence of unexpected results as shown by the data provided in the originally filed specification provide sufficient basis for demonstration of a surprising and synergistic effect provided by the combination of ST1326 and metformin - even at subpharmaceutical doses. The results presented in the original specification accompanied by the executed declaration signed by the inventors would have significant evidentiary weight, comparable to the weight given to an executed declaration. It is well established by the Federal Circuit that "the examiner must consider comparative data presented in the specification which is intended to illustrate the claimed invention in reaching a conclusion in regard to the obviousness of claims." In re Margolis, 785 F.2d 1029, 228 U.S.P.Q. 1123, 1129 (Fed. Cir. 1993).

For the above reasons it is respectfully submitted that claims 1 and 9-17 define inventive subject matter and that claims 3-8 should be rejoined and all pending claims allowed.

Respectfully submitted,

NIXON & VANDERHYE P.C.

By:

Arthur R. Crawford Reg./No. 25,327

ARC:eaw

901 North Glebe Road, 11th Floor

Arlington, VA 22203-1808

Telephone: (703) 816-4000

Facsimile: (703) 816-4100